INTENDED USE
The Spectrum H. pylori Ab Test Device is a sandwich lateral flow chromatographic immunoassay for the qualitative detection of antibodies (IgG, IgM, and IgA) anti-Helicobacter pylori (H. pylori) in human serum, plasma, whole blood. It is intended to be used as a screening test and as an aid in the diagnosis of infection with H. pylori. Any reactive specimen with the Spectrum H. pylori Ab Test Device must be confirmed with alternative testing method(s) and clinical findings.

SUMMARY AND EXPLANATION OF THE TEST
Helicobacter pylori is associated with a variety of gastrointestinal diseases including non-ulcer dyspepsia, duodenal and gastric ulcer and active, chronic gastritis. The prevalence of H. pylori infection could exceed 90% in patients with signs and symptoms of gastrointestinal diseases. Recent studies indicate an association of H. pylori infection with stomach cancer.

H. pylori colonizing in the gastrointestinal system elicits specific antibody responses which aids in the diagnosis of H. pylori infection and in monitoring the prognosis of the treatment of H. pylori related diseases. Antibiotics in combination with bismuth compounds have been shown to be effective in treating active H. pylori infection. Successful eradication of H. pylori is associated with clinical improvement in patients with gastrointestinal diseases providing a further evidence.

The Spectrum H. pylori Ab Test Device is a latest generation of chromatographic immunotest which utilizes recombinant antigens to detect the antibodies to H. pylori in human serum or plasma. The test is user friendly, highly sensitive and specific.

TEST PRINCIPLE
The Spectrum H. pylori Ab Test Device is a lateral flow chromatographic immunotest based on the principle of the double antigen–sandwich technique. The test cassette consists of:1) a burgundy colored conjugate pad containing H. pylori antigens including Cag-A conjugated with colloidal gold (H. pylori conjugates) and rabbit IgG-gold conjugates, 2) a nitrocellulose membrane strip containing a test band (T band) and a control band (C band). The T band is pre-coated with non-conjugated H. pylori antigens, and the C band is pre-coated with goat anti-rabbit IgG.

When an adequate volume of test specimen is dispensed into the sample well of the cassette, the specimen migrates by capillary action across the cassette. The antibodies: either the IgG, the IgM, or the IgA, to H. pylori if present in the specimen will bind to the H. pylori conjugates. The immunocomplex is then captured on the membrane by the pre-coated H. pylori antigens, forming a burgundy colored T band, indicating a H. pylori Ab positive test result.

Absence of the T band suggests a negative result. The test contains an internal control (C band) which should exhibit a burgundy colored band of the immunocomplex of goat anti-rabbit IgG/rabbit IgG-gold conjugate regardless the presence of any antibodies to H. pylori. Otherwise, the test result is invalid and the specimen must be retested with another device.

REAGENTS AND MATERIALS PROVIDED
1. Individually sealed foil pouches containing:
   a. One cassette device.
   b. One plastic dropper.
   c. One desiccant.
2. Sample Diluent (1 bottle, 5 mL)
3. One package insert (instruction for use).

MATERIALS MAY BE REQUIRED AND NOT PROVIDED
1. Positive Control
2. Negative Control

MATERIALS REQUIRED BUT NOT PROVIDED
1. Clock or Timer
2. Lancing device for whole blood test

WARNINGS AND PRECAUTIONS

For in Vitro Diagnostic Use
1. This package insert must be read completely before performing the test. Failure to follow the insert gives inaccurate test results.
2. Do not open the sealed pouch, unless ready to conduct the assay.
3. Do not use expired devices.
4. Bring all reagents to room temperature (15°C-30°C) before use.
5. Do not use the components in any other type of test kit as a substitute for the components in this kit.
6. Do not use hemolized blood specimen for testing.
7. Wear protective clothing and disposable gloves while handling the kit reagents and clinical specimens. Wash hands thoroughly after performing the test.
8. Users of this test should follow the US CDC Universal Precautions for prevention of transmission of HIV, HBV and other blood-borne pathogens.
9. Do not smoke, drink, or eat in areas where specimens or kit reagents are being handled.
10. Dispose of all specimens and materials used to perform the test as biohazardous waste.
11. Handle the Negative and Positive Control in the same manner as patient specimens.
12. The testing results should be read within 15 minutes after a specimen is applied to the sample well or sample pad of the device. Read result after 15 minutes may give erroneous results.
13. Do not perform the test in a room with strong air flow, ie. electric fan or strong air-conditioning.

REAGENT PREPARATION AND STORAGE INSTRUCTIONS
All reagents are ready to use as supplied. Store unused test device openened at 2°C-30°C. If stored at 2°C-8°C, ensure that the test device is brought to room temperature before opening. The test device is stable through the expiration date printed on the sealed pouch. Do not freeze the kit or expose the kit over 30°C.

SPECIMEN COLLECTION AND HANDLING
Consider any materials of human origin as infectious and handle them using standard biosafety procedures.

Plasma
1. Collect blood specimen into a lavender, blue or green top collection tube (containing EDTA, citrate or heparpin, respectively in Vacutainer®) by veinpuncture.
2. Separate the plasma by centrifugation.
3. Carefully withdraw the plasma into new pre-labeled tube.
**ASSAY PROCEDURE**

**Step 1:** Bring the specimen and test components to room temperature if refrigerated or frozen. Mix the specimen well prior to testing. Place the test device on a clean, flat surface.

**Step 2:** When ready to test, open the pouch at the notch and remove the device. Be sure to label the device with specimen’s ID number.

**Step 3:** Fill the plastic dropper with the specimen. Holding the dropper vertically, dispense 1 drop (about 30-45 µL) of serum/plasma or 1 drop of whole blood (about 40-50 µL) into the sample well making sure that there are no air bubbles. Immediately add 1 drop (about 35-50 µL) of Sample Diluent to the sample well.

**Step 5:** Set up timer. Results can be read in 15 minutes. Positive results can be visible in as short as 1 minute.

**Step 6:** Samples with positive results should be confirmed with alternative testing method(s) and clinical findings before a positive determination is made.

**PERFORMANCE CHARACTERISTICS**

**Clinical Performance**

A total of 200 specimens from the non- H. pylori patients and 75 specimens from the patients under anti-H. pylori treatment were tested by the Spectrum H. pylori Ab Test Device. Comparison for all subjects is shown in the following table.

<table>
<thead>
<tr>
<th>Spectrum H. pylori Ab Test Device</th>
<th>H. pylori Patients</th>
<th>Positive</th>
<th>Negative</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive</td>
<td>65</td>
<td>10</td>
<td>75</td>
<td></td>
</tr>
<tr>
<td>Negative</td>
<td>18</td>
<td>182</td>
<td>200</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>83</td>
<td>180</td>
<td>275</td>
<td></td>
</tr>
</tbody>
</table>

Relative Sensitivity: 86.7%, Relative Specificity: 91%, Overall Agreement: 89.8%

**LIMITATIONS OF TEST**

1. The Assay Procedure and the Assay Result Interpretation must be followed closely when testing the presence of antibodies to H. pylori in serum, plasma or whole blood from individual subjects. Failure to follow the procedure may give inaccurate results.

2. The Spectrum H. pylori Ab Test Device is limited to the qualitative detection of IgG, IgM, and IgA anti- H. pylori in human serum, plasma or whole blood. The intensity of the test band does not have linear correlation with the antibody titer in the specimen.

3. A negative result for an individual subject indicates absence of detectable antibodies to H. pylori. However, a negative test result does not preclude the possibility of exposure to or infection with H. pylori.

4. A negative result can occur if the quantity of the antibodies to H. pylori present in the specimen is below the detection limits of the assay, or the antibodies that are detected are not present during the stage of disease in which a sample is collected.

5. Some specimens containing unusually high titer of heterophile antibodies or rheumatoid factor may affect expected results.

6. The results obtained with this test should only be interpreted in conjunction with other diagnostic procedures and clinical findings.

**REFERENCES**


**QUALITY CONTROL**

1. Internal Control: This test contains a built-in control feature, the C band. The C line develops after adding specimen and sample diluent. Otherwise, review the whole procedure and repeat test with a new device.

2. External Control: Good Laboratory Practice recommends using the external controls, positive and negative, to assure the proper performance of the assay, particularly under the following circumstancs:
   a. A new operator uses the kit, prior to performing testing of specimens.
   b. A new kit is used.
   c. A new shipment of kits is used.
   d. The temperature used during storage of the kit falls outside of 2-15°C.
   e. The temperature of the test area falls outside of 15 - 30°C.
   f. To verify a higher than expected frequency of positive or negative results.
   g. To investigate the cause of repeated invalid results.